

Article



Restricted Geographic Sampling Yields Low Parasitism Rates but Surprisingly Diverse Host Associations in Avian Lice (Insecta: Phthiraptera) from South Texas

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Abstract: South Texas is a highly variable region encompassing multiple habitat types and harboring a wide diversity of organisms. However, the parasite fauna in this region is poorly known, especially for avian ectoparasites such as lice. To better understand avian louse diversity and host associations in South Texas, we examined a total of 507 birds for chewing lice. Lice were morphologically identified to genus and phylogenetic analysis was performed using one mitochondrial (COI) and two nuclear (18S rRNA and EF-1 α) genes. Of the birds examined, 69 (13.5%) were parasitized by lice resulting in a total of 63 host associations across 45 bird species, 29 of which were previously unrecorded. The predominant taxa encountered during this study included two of the most diverse louse genera, *Myrsidea* and *Brueelia*. Molecular analyses revealed 21 distinct genetic lineages, 17 of which are associated with novel host associations and may represent new species. This study represents the first extensive examination of avian louse host associations and relationships in Texas and reveals that there is still much to be learned about ectoparasite diversity in the New World.

Keywords: avian lice; birds; chewing lice; ectoparasites; genetics; parasitism

1. Introduction

Chewing lice (Insecta: Phthiraptera) are small, dorsoventrally flattened ectoparasites found on numerous bird and mammal species [1,2]. Avian chewing lice are placed into two suborders, Amblycera and Ischnocera. Ischnocerans are feather specialists (feather lice) feeding on feathers and dead skin. Amblycerans have a similar diet, but some also feed on blood or body fluid secretions [3]. Each suborder is characterized by distinctive morphologies, and they partition the avian body based on feeding strategy and host preening avoidance [4]. Lice are obligate, permanent parasites that spend their entire life cycle (egg, nymphs, and adults) on their host (i.e., there is no free-living stage to the life cycle [5]). However, chewing lice can move among hosts via two main methods of transport: vertical transmission from parent to offspring, and horizontal transmission via phoresis ("hitchhiking") on hippoboscid flies [1,6,7]. Although phoresis is not uncommon, it is more frequently used by ischnoceran than amblyceran lice [1,7,8]. This method of transmission may allow lice to escape competition from other lice on the same host and potentially encounter a novel host [8]. Despite the ability to switch hosts (should the opportunity occur), many louse species are highly specialized to one host species [2]. However, some louse species and genera can be host generalists, parasitizing multiple host species over a large geographic area. Sometimes these host generalists may represent cryptic species [9], but many times these lice simply have wide host and geographic associations. Close morphological examinations as well as genetic assessments of lice are needed to better



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Copyright: © 2021 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). understand host associations, louse relationships, and to determine if generalist species are harboring cryptic species or simply have a widespread host and geographic range.

Several studies have investigated the louse associations of avian species in North America. Exemplar studies include examination of ectoparasites from Wild Turkey (*Meleagris gallopavo*) in California [10], community structure of lice on the Western Scrub Jay (*Aphelocoma californica*) in the Southwestern United States [11], phylogenetic relationships of lice on *Catharus* thrushes in Illinois [12], and louse–host associations of dove species in Manitoba, Canada [13]. Except for a few studies examining population genetics, parasitism rates, and host switching of chewing lice parasitizing doves [14–16], there have been few studies that have examined louse relationships in Texas. This is surprising as Texas and especially South Texas, is highly diverse in terms of habitats and vertebrate fauna [17]. As such, Texas is not unlike Mexico and Central America in being an understudied area with respect to understanding louse–host associations. For example, several studies in Mexico have examined co-evolutionary processes and speciation of chewing lice parasitizing birds, but these have been limited to dove species [9,14,18]. In Central America, only a few studies from Costa Rica [19–21] and Panama [22] exist, but these primarily provide descriptions of new louse species rather than descriptions of host associations and louse diversity.

Given these limited studies, many questions remain unanswered about the biogeography, host associations, and genetic relationships of avian chewing lice across a significant portion of North America. The objectives of this study are to assess louse–host associations and louse diversity, and to construct molecular phylogenies to determine relationships among lice collected from birds surveyed in South Texas.

2. Materials and Methods

2.1. Louse Sampling and Identification

Chewing lice were collected from birds intermittently during 2013–2015 on two East Foundation (eastfoundation.net) ranches located in southern Texas (Figure 1). These East Foundation properties occupy approximately 71,200 hectares of land in Jim Hogg, Starr, Willacy, and Kenedy Counties. For this study, samples were obtained from San Antonio Viejo Ranch (60,298 hectares located inland in Jim Hogg and Starr Counties) and El Sauz Ranch (10,926 hectares located coastally in Kenedy and Willacy Counties; Figure 1). While the two ranches are generally similar in terms of dominant vegetation, El Sauz has extensive sand sheets, ephemeral ponds, and receives ca. 10 cm more rain annually (30-year Normals via [23]) as compared to San Antonio Viejo. The avifaunal composition of the two ranches is effectively the same in terms of resident and wintering species, although El Sauz appears to have a greater number of Neotropical migrants passing through, likely due to its coastal location (JPP pers. obs.).

Birds were captured in the field via mist net during all seasons. Some birds were released, and others were euthanized and prepared as voucher specimens and deposited in either the East Foundation's collection or at the Biodiversity Research and Teaching Collections (BRTC) located at Texas A&M University (TAMU; according to TAMU Animal Care and Use Permits IACUC 2012-6 and 2015–0020). In both instances, birds were ruffled for chewing lice using a toothbrush to thoroughly brush the feathers over collecting paper, with a new piece of paper being used for each bird [24,25]. All ectoparasites were transferred to a collection tube. The birds that were released were dusted with flea powder (Zodiac[®], Schaumburg, IL, USA) to aid in releasing ectoparasites from their plumage; birds were ruffled prior to release [26]. Each euthanized bird was placed in its own individual bag with a cotton ball dipped in ethyl acetate, which acts as a fumigant to release the lice from their host [25]. After several minutes of exposure to ethyl acetate, birds were ruffled, and lice transferred to a collection tube as above. After lice were collected from a bird, they were stored either dry or in ethanol at -80 °C.



Figure 1. Map of Texas, with an emphasis on south Texas and all East Foundation properties (shaded regions). The two properties for this study (San Antonio Viejo and El Sauz) are indicated.

Digital vouchers were created before DNA extraction (below) for each chewing louse by using an Olympus SZX10 microscope, Intralux 6000 light source and SPOT v4.6 software (2009 Diagnostic Instruments). Following DNA extraction, specimen exoskeletons were retained, cleared, and mounted on a microslide in balsam as vouchers, following general protocols [27]. We used a combination of published keys, taxonomic descriptions, and previously identified vouchers with molecular data to identify our specimens to the lowest taxonomic level possible.

We used Price et al. [2] to identify our specimens to genus and to determine chewing louse species that were previously recorded from the host species in question. We elected not to make species determinations based solely on published host records. This is partially due to the fact many hosts are poorly sampled, particularly in the geographic region we focused on. It is also possible that multiple congeneric lineages of lice exist on a single host species in different portions of the range [28] leading to erroneous identifications. When possible, we used louse species identification keys, but in most instances, we relied primarily on original descriptions and revisions. Specifically, we utilized the works of Carriker [29], Clay [30,31], Dalgleish [32], Emerson [33,34], Grossi et al. [35], Kellogg [36], Osborn [37], Price [38,39], and Williams [40] to identify our specimens. Although we attempted to identify every voucher, in some instances, our specimens were not able to be identified with confidence. This included instances where nymphs were sequenced or if the voucher was missing structures required for identification. In some cases, multiple samples representing the same louse species were sequenced. In these cases, other vouchers were identifiable, and that identification was applied across all samples. In a few instances, published papers included species level identification of their sequenced louse samples. In instances where our sequence data indicated we had conspecific lice in our sample, we included these identifications. In the remaining instances, we left the species identification undetermined, regardless of if there was a published host

association between the louse genus and the host species. No identification was made based exclusively on host-parasite relationship.

2.2. DNA Extraction, Amplification, and Sequencing

DNA extraction was performed using the E.Z.N.A.[®] Tissue DNA Kit (Omega Bio-Tek Inc., Norcross, GA, USA) following louse specific protocols [41]. Before the extraction began, the chewing lice were washed in 1X phosphate-buffered saline solution to remove possible contaminants from the specimen. After the wash, the abdomen of the louse specimen was sliced using a sterile surgical blade. The manufacturer's extraction protocol was followed throughout the process, except the total DNA elution volume was lowered to 70 μ L (35 μ L per elution). Upon completion of DNA extraction, each louse exoskeleton was preserved in ethanol and stored at -80° C to be retained as a voucher specimen at the Texas A&M University Insect Collection.

Polymerase chain reaction (PCR) was performed to amplify a portion of the mitochondrial cytochrome c oxidase subunit I (COI) gene, as well as a portion of the nuclear 18S ribosomal RNA (18S) and elongation factor-1 α (EF-1 α) genes. We used primers L6625 and H7005 to isolate COI [42], NS1 and NS2a for 18S [43], and EF1-For3 and Cho10 for EF-1 α [44]. Amplification protocols for COI and EF-1 α genes were followed as described in Light et al. [45] and for the 18S protocol as described in Light et al. [46]. We amplified COI for all samples from which DNA was isolated, while amplifications of 18S and EF-1 α were performed for a subset of taxa that represented individual clades recovered in the COI phylogenies. The amplified PCR products were electrophoresed with 2 µL of 100 bp Promega DNA ladder (Applied Biosystems) on an agarose gel to determine PCR success. Successfully amplified PCR products were purified using ExoSAP-IT (United States Biochemical Corporation, Cleveland, OH, USA) and sent to Beckman-Coulter Genomics (Danvers, MD, USA) or Yale University (New Haven, CT, USA) for sequencing in both forward and reverse directions using the PCR primers. Forward and reverse strands were edited and combined using Sequencher v.4.2.2 (Gene Codes Corporation, Ann Arbor, MI, USA). Sequences were then aligned using MUSCLE [47] and verified by eye. Sequences are available on GenBank (Table S2).

2.3. Sequence Analyses

Each chewing louse sequence was compared to published sequences located in GenBank by using the Basic Local Alignment Search Tool (BLAST). Closely related sequences resulting from the BLAST search (Table S1) were added to our data set for phylogenetic analyses. Phylogenetic analysis for COI occurred in two separate analyses: one for the suborder Amblycera and one for the suborder Ischnocera. Each analysis used two specimens from the louse suborder Anoplura as outgroups (Fahrenholzia zacatecae and Haematopinus eurysterunus; Table S1). Analysis of the nuclear genes, 18S and EF-1 α , was performed for each gene separately but with Amblycera and Ischnocera combined in each analysis. In addition to these individual gene analyses, four different analyses were performed combining genes to assess louse relationships: two using mitochondrial and nuclear genes (COI+18S, COI+EF-1α), one assessing only the nuclear genes ($18S+EF-1\alpha$), and one analysis combining all genes $(COI+18S+EF-1\alpha)$ using concatenated data sets. Phylogenetic analyses of these combinations of genes were performed to illustrate how the more rapidly evolving COI gene was driving tree topology. Amblycera and Ischnocera were combined in each of these four combined gene analyses. Although we were not able to amplify all genes for all taxa and many GenBank specimens did not have all three genes available (Tables S1 and S2), we included all specimens and their available data in each phylogenetic analysis.

Prior to phylogenetic analysis, PartitionFinder v2.1.1 [48] was used with the Bayesian information criterion to select the appropriate number of partitions and model of evolution at each partition for each gene. The same partitions and models of evolution identified in the single gene data sets were used in the combined gene analyses. For both protein coding genes, three partitions were selected, one for each codon position (COI: HYK+G,

GTR+I+G, and GTR+I+G; EF-1 α : K80+I+G, SYM+I+G, and K80+I+G, for the first, second, and third codon positions, respectively), whereas only one partition was selected for 18S (SYM+I+G). Bayesian phylogenetic analyses were performed for each individual gene and combined gene data sets using MrBayes 3.2 [49]. Analyses for each data set consisted of two simultaneous runs for 10 million generations with four heated chains [50] and sampling occurred every 2000 generations with a 25% burn-in. Tracer v1.6 [51] was used to assess convergence of the data. A 50% majority rule consensus tree was constructed in FigTree v1.4.2 [52] for each analysis and the percentage of samples recovered in a particular clade was assumed to represent that clade's posterior probability [53]. Average uncorrected *p*-distances were calculated for COI in PAUP* v4.0 [54] to examine the genetic difference among taxa. Genetic lineages were determined to be potentially distinct if they differed by a minimum of 15% (uncorrected p-distance) from their closest relative. Statistical analysis was performed using the Fisher's exact test (*p* < 0.05) to assess louse parasitism from each ranch and was performed in R [55].

3. Results

3.1. Parasitism Rates

A total of 507 bird specimens from two sampling localities in South Texas were examined for chewing lice. The host specimens examined were taxonomically diverse, representing a total of 140 species, 36 families, and 12 orders (Table S3). Of these taxonomic categories, 32.1%, 58.3%, and 75.0%, were parasitized by ectoparasites, respectively. There were 221 birds collected from San Antonio Viejo Ranch (10.4% were parasitized) and 286 birds collected from El Sauz (16.1% parasitized; Table 1), although this difference in parasitism rates was not statistically significant (Fisher's exact test, p = 0.0685). In total, 69 birds were parasitized by lice (13.5%) and 63 host associations were recorded, with 29 of these associations being new (Table 2). There were 12 instances in which a host species was parasitized by more than one species of louse (Table 2). Lice from both suborders were found to be parasitizing the same host species eight times (Table 2). We note that some lice were not identified to species due to lack of reference material. Although not all specimens could be identified to species, species identity could sometimes be determined based on genetic similarity to specimens on GenBank. Additional host associations may also have been present in this study, but some louse specimens were excluded due to poor condition of the specimens or lack of reference material.

Host Order	Host Family	Host Species (n)	Locality
Anseriformes	Anatidae	Northern Shoveler (Anas clypeata) (1)	El Sauz
Galliformes	Odontophoridae	Northern Bobwhite (Colinus virginianus) (1)	El Sauz
Accipitriformes	Accipitridae	Harris's Hawk (Parabuteo unicinctus) (1)	El Sauz
Falconiformes	Falconidae	American Kestrel (Falco sparverius) (1)	San Antonio Viejo
Charadriformes	Scolopacidae	Sanderling (<i>Calidris alba</i>) (1)	El Sauz
	1	Long-billed Curlew (Numenius americanus) (1)	El Sauz
Columbiformes	Columbidae	Common Ground Dove (<i>Columbina passerina</i>) (1)	El Sauz
		Mourning Dove (Zenaida macroura) (1)	El Sauz
Cuculiformes	Cuculidae	Yellow-billed Cuckoo (<i>Coccuzus americanus</i>) (1)	San Antonio Vieio
Piciformes	Picidae	Golden-fronted Woodpecker (Melanerpes aurifrons) (1)	San Antonio Viejo
Passeriformes	Alaudidae	Horned Lark (Eremophila alvestris) (1)	El Sauz
	Cardinalidae	Northern Cardinal (<i>Cardinalis cardinalis</i>) (6)	El Sauz
		Northern Cardinal (<i>Cardinalis cardinalis</i>) (1)	San Antonio Vieio
		Painted Bunting (Passerina ciris) (1)	San Antonio Vieio
		Varied Bunting (Passerina versicolor) (1)	San Antonio Vieio
		Summer Tanager (<i>Piranga rubra</i>) (2)	El Sauz
	Corvidae	Green Jay (<i>Cyanocorax yncas</i>) (2)	El Sauz
	Emberizidae	Lark Sparrow (Chondestes grammacus) (1)	El Sauz
		Lincoln's Sparrow (Melospiza lincolnii) (1)	El Sauz
		Cassin's Sparrow (Pecaea cassini) (1)	San Antonio Vieio
	Fringillidae	Lesser Goldfinch (<i>Spinus psaltria</i>) (1)	El Sauz
	Iceteridae	Red-winged Blackbird (Agelaius phoeniceus) (4)	San Antonio Vieio
	recontrace	Hooded Oriole (Icterus cucullatus) (1)	El Sauz
		Audubon's Oriole (Icterus graduacauda) (1)	San Antonio Vieio
		Bronzed Cowbird (Molothrus aeneaus) (2)	El Sauz
		Bronzed Cowbird (Molothrus aeneaus) (1)	San Antonio Vieio
		Brown-headed Cowbird (<i>Molothrus ater</i>) (1)	El Sauz
		Eastern Meadowlark (Sturnella magna) (1)	San Antonio Vieio
	Mimidae	Northern Mockingbird (<i>Minus polyglottos</i>) (6)	El Sauz
		Northern Mockingbird (<i>Minus polygiotics</i>) (1)	San Antonio Vieio
		Curve-billed Thrasher (Toxostoma curvirostre) (2)	San Antonio Viejo
		Long-billed Thrasher (Toxostoma longitostre) (2)	El Sauz
		Long-billed Thrasher (Torostoma longitostre) (1)	San Antonio Vieio
	Hirundinidae	Tree Swallow (<i>Tachycineta hicolor</i>) (1)	El Sauz
	Parulidae	Black-and-white Warbler (<i>Mniotilta varia</i>) (1)	El Sauz
	1 ur ur ur ur u	Tennessee Warbler (Oreothlynis peregrina) (1)	San Antonio Vieio
		Northern Waterthrush (Parkesia noeboracensis) (1)	El Sauz
		Louisiana Waterthrush (Parkesia motacilla) (2)	El Sauz
		Northern Parula (Setonhaga americana) (1)	El Sauz
		Bay-breasted Warbler (Setonhaga castanea) (1)	El Sauz
		Black-throated Green Warbler (Setonhaga virens) (1)	El Sauz
		Golden-winged Warbler (Vermivora chrysontera) (1)	El Sauz
	Remizidae	Verdin (Auriparus flavicens) (1)	San Antonio Vieio
	Troglodvtidae	Cactus Wren (<i>Campylorhynchus brunneicavillus</i>) (1)	San Antonio Vieio
	110910ay trade	Carolina Wren (<i>Thruothorus ludovicianus</i>) (1)	El Sauz
	Tyrannidae	Brown-crested Flycatcher (<i>Mujarchus turannulus</i>) (1)	El Sauz
	1) Infinitude	Vermilion Flycatcher (<i>Purocenhalus ruhinus</i>) (1)	San Antonio Vieio
		Scissor-tailed Tlycatcher (Turannus forficatus) (1)	El Sauz
	Vireonidae	White-eved Vireo (Vireo oriseus) (1)	El Sauz

Table 1. Species of birds that were parasitized by chewing lice in South Texas from two East Foundation ranches (San Antonio Viejo and El Sauz) during 2013–2015, where (n) is total number of host individuals per species examined.

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Host Order	Host Family	Host Species	Louse Suborder	Louse Species
Anseriformes	Anatidae	Anas cluneata	Ischnocera	Anaticola crassicornis
ritsernormes	7 mandae	111105 cryptutu	Ischnocera	Anatoecus sp
Galliformes	Odontophoridae	Colinus viroinianus	Ischnocera	Goniodes sp. *
Accipitriformes	Accipitridae	Parabuteo unicinctus	Ischnocera	Degeeriella fulva †
Falconiformes	Falconidae	Falco sparverious	Ischnocera	Degeeriella sp.
Charadriformes	Scolopacidae	Calidris alba	Ischnocera	Lunaceps actophilus
	1		X 1	Cummingsiella
		Numenius americanus	Ischnocera	longirostricola
Columbiformes	Columbidae	Columbina passerina	Ischnocera	Columbicola passerinae
		,	Amblycera	Hohorstiella passerinae
		Zenaida macroura	Ischnocera	Columbicola macrourae
Cuculiformes	Cuculidae	Coccyzus americanus	Ischnocera	<i>Cuculoecus</i> sp. *
Piciformes	Picidae	Melanerpes aurifrons	Ischnocera	Penenirmus sp. *
			Ischnocera	Picicola snodgrassi
Passeriformes	Alaudidae	Eremophila alprestris	Amblycera	<i>Menacanthus</i> sp. †*
	Cardinalidae	Cardinalis cardinalis	Ischnocera	Brueelia pallidula +
			Amblycera	Menacanthus
			Timorycera	eurysternus
			Amblycera	<i>Myrsidea</i> sp. †
		Passerina ciris	Amblycera	<i>Myrsidea</i> sp. +*
			Ischnocera	Philopterus sp. +*
		Passerina versicolor	Amblycera	<i>Ricinus</i> sp.
		Piranga rubra	Amblycera	<i>Myrsidea</i> sp. +*
		_	Ischnocera	Philopterus sp. +*
	Corvidae	Cyanocorax yncas	Amblycera	Menacanthus sp.
			Amblycera	<i>Myrsidea</i> sp. †*
			Ischnocera	Philopterus sp. †*
	Emberizidae	Chondestes grammacus	Amblycera	<i>Myrsidea</i> sp. †*
		Melospiza lincolnii	Ischnocera	Brueelia sp.
		Peucaea cassinii	Ischnocera	Philopterus sp. †*
	Fringillidae	Spinus psaitria	Ischnocera	Philopterus sp. T
	Hirundinidae	lachycineta bicolor	Amblycera	Myrsiaea sp. †*
	Iceteridae	Agenaus phoeniceus	Ischnocera	Brueella xanthocephali
			Amblycera	Murcidea op +*
			Ischnosera	Philopterus sp. *
		Ictory cucullatur	Amblycora	Manacanthus sp. +
		icierus cuculturus	Ischnocera	Philopterus sp. +*
		Icterus oraduacauda	Ischnocera	Brueelia miloata
		Molothrus aeneaus	Ischnocera	Brueelia xanthocenhali
		Molothrus ater	Ischnocera	Brueelia xanthocephali
		Woldthin us aler	Amblycera	Mursidea sp
		Sturnella maona	Ischnocera	Brueelia nicturata
	Mimidae	Mimus polyglottos	Ischnocera	Brueelia brunneinucha
			Amblycera	Mursidea sp. †*
			Ischnocera	Philopterus sp. *
		Toxostoma curvirostre	Ischnocera	Brueelia dorsale
		Toxostoma longirostre	Ischnocera	Brueelia brunneinucha +
		8	Ischnocera	Brueelia dorsale +
			Ischnocera	Philopterus sp. +*
			Ischnocera	Brueelia sp. + *
	Parulidae	Mniotilta varia	Ischnocera	Philopterus sp. †*
		Oreothlypis peregrina	Amblycera	<i>Ricinus</i> sp. *
		Parkesia noveboracensis	Amblycera	Ricinus sp. *
		Parkesia motacilla	Amblycera	<i>Myrsidea</i> sp. *
		Setophaga americana	Amblycera	Menacanthus sp. +*
		Setophaga virens	Amblycera	<i>Myrsidea</i> sp. †
		Vermivora chrysoptera	Amblycera	<i>Ricinus</i> sp. *

Table 2. Louse–host associations of birds sampled from South Texas during 2013–2015. Daggers (†) indicate new host associations and asterisks (*) indicate species with no molecular data.

Host Order	Host Family	Host Species	Louse Suborder	Louse Species
	Remizidae	Auriparus flaviceps	Ischnocera	Brueelia sp. *
	Troglodytidae	Campylorhynchus brunneicapillus	Ischnocera	Brueelia dorsale +
		Thryothorus ludovicianus	Amblycera	<i>Myrsidea</i> sp.
	Tyrannidae	Myiarchus tyrannulus	Amblycera	Menacanthus sp. +*
		Pyrocephalus rubinus	Amblycera	<i>Ricinus</i> sp.
		Tyrannus forficatus	Ischnocera	Picicola sp. *
	Vireonidae	Vireo griseus	Ischnocera	Brueelia dorsale +

Table 2. Cont.

3.2. COI Analyses

There were a total of 148 chewing louse specimens included in the COI phylogenetic analysis: 81 specimens (35 Amblycera and 46 Ischnocera) from South Texas and 67 (20 Amblycera, 45 Ischnocera, and two Anoplura) from GenBank (Figures 2 and 3). There were high average pairwise sequence divergences (uncorrected *p*-distances) within each suborder: across amblyceran taxa, the average uncorrected *p*-distance was 23.3% with a range of 0–35.7%; across ischnoceran taxa, the average uncorrected *p*-distance was 26.1% with a range of 0–37.3%. From this study, there were 21 distinct genetic lineages (seven from Amblycera and 14 from Ischnocera), each of which was approximately 15% genetically divergent (uncorrected *p*-distance) from their closest relative (Figures 2 and 3).

Phylogenetic analysis of the amblyceran taxa resulted in three clades with strong Bayesian posterior probability (PP) support; each clade circumscribes a major louse genus (Figure 2). *Myrsidea* had the strongest support (PP = 1), and the *Ricinus* and *Menacanthus* clades were strongly supported as well (PP = 0.99). Support for relationships within each clade varied. In *Menacanthus* and *Ricinus*, more relationships were strongly supported when compared to *Myrsidea*. Within *Myrsidea*, most relationships were poorly supported, effectively resulting in a major polytomy within this genus. The analysis also included the amblyceran genus *Hohorstiella*, which was placed as sister to *Menacanthus*; however, this relationship was poorly supported (PP = 0.73).

Analysis of the ischnoceran suborder revealed several well-supported clades (Figure 3). The louse genera *Columbicola, Anaticola*, and *Anatoecus* were recovered together in a clade, with posterior support of 0.94. The *Degeeriella* complex, comprising the genera *Degeeriella*, *Picicola, Trogoninirmus*, and *Contingacola*, was strongly supported (PP = 1), although intergeneric relationships were not strongly supported and *Picicola* was not recovered as monophyletic. Members of the louse genera *Cummingsiella, Lunaceps, Penenirmus* (not recovered as monophyletic), *Cuculoecus*, and *Goniodes* comprised a poorly supported clade, sister to *Philopterus* (PP = 0.99). While *Philopterus* was strongly supported (PP = 0.99), the relationships within the genus were not. The same is true for the largest ischnoceran clade that comprised of members of the hyper-diverse genus *Brueelia*. This clade was strongly supported (PP = 1); however, support for relationships within the clade was lacking.

3.3. 18S, EF-1 α , and Combined Analyses

A total of 49 chewing louse specimens were included in the 18S phylogenetic analyses: 44 specimens from South Texas (16 Amblycera and 28 Ischnocera) and five from GenBank (one Amblycera, two Ischnocera, and two Anoplura; Figure S1). There were 55 specimens included in the EF-1 α analysis, 36 lice from South Texas (16 Amblycera, 20 Ischnocera) and 19 from GenBank (six Amblycera, nine Ischnocera and two Anoplura; Figure S2). The large clades recovered in the COI analysis were recovered in both 18S and EF-1 α . A monophyletic Ischnocera was not supported with analysis of EF-1 α (Figure S2). Each gene analysis revealed slightly different relationships within those large clades, although they were poorly supported.



Figure 2. Bayesian phylogram of amblyceran taxa from south Texas. Posterior probabilities ≥ 0.95 are shown and sequences in gray are from GenBank. Distinct genetic lineages are denoted with asterisks (*); one of these lineages (*Menacanthus distinctus*) represents a morphologically described taxon. Taxon labels for newly sequenced specimens include the louse species (if identified), unique identifier (generally matching the host voucher number), and host species. Taxon labels for GenBank specimens include louse species (if known), host species, and sampling location (if known). Outgroup taxa have been pruned.



Figure 3. Bayesian phylogram of ischnoceran taxa from south Texas. Posterior probabilities \geq 0.95 are shown and sequences in gray are from GenBank. Distinct genetic lineages are denoted with asterisks (*); two of these lineages (*Goniodes ortygis, Penenirmus auritus*) represent morphologically described taxa. The host families Icteridae and Mimidae are shown to the right of louse clades exclusively shown here to parasitize them. Taxon labels for newly sequenced specimens include the louse species (if identified), unique identifier (generally matching the host voucher number), and host species. Taxon labels for GenBank specimens include louse species (if known), host species, and sampling location (if known). Outgroup taxa have been pruned.

When the two nuclear genes, 18S and EF-1 α (52 samples from South Texas and 16 from GenBank), were analyzed together, the results gave high support for the suborders and many of the genera and intergeneric relationships (Figure S3). The analysis of COI+18S (45 samples from South Texas and three from GenBank) provided resolved and well-supported relationships (Figure S4), more so than the analysis of the combined nuclear genes (Figure S3). The relationships seen in the tree resulting from the COI+EF-1 α analysis (37 samples from South Texas and 16 from GenBank) also received high support, although Ischnocera

was again not supported as monophyletic (Figure S5). When all three genes were combined into one analysis (50 samples from South Texas and 18 from GenBank), suborder, genera, and intergeneric relationships were resolved and well-supported, more so than any of the analyses of individual or combined gene data sets (Figure 4). Notably, this three-gene tree topology is very similar to that seen in the COI-only trees (Figures 2 and 3), although relationships were better resolved and supported across the tree (Figure 4).



Figure 4. Bayesian phylogram of combined mitochondrial (COI gene) and nuclear genes (18S and EF-1 α). Posterior probabilities ≥ 0.95 are shown and sequences in gray are from GenBank. Taxon labels for newly sequenced specimens include the louse species (if identified), unique identifier (generally matching the host voucher number), and host species. Taxon labels for GenBank specimens include louse species (if known) and host species. Outgroup taxa have been pruned.

4. Discussion

With respect to parasitism rates, the bird species examined for chewing lice in this study represent a taxonomically diverse subset of species occurring in South Texas. Overall, we recovered a parasitism rate of 13.5%, despite having sampled across seasons. Indeed, our recovered parasitism rate would have been even lower, had we not sampled migratory birds in spring (e.g., Parulidae taxa). Previous studies that have assessed broad assemblages of host species for lice have found somewhat similar parasitism rates of 8.3% (Benin [56]), 19% (Democratic Republic of the Congo [45]), 20% (Brazil [57]), and 22% (South Africa [58]). The findings of this study were not consistent with the extraordinarily high parasitism rate of 60% or more observed from Central America, South America, and Europe [19,59–61]. Some of these studies reflect parasitism rates from a single season sampling event (e.g., [58]), while others reflect parasitism rates from across multiple seasons (e.g., [57]).

While chewing louse prevalence may vary geographically or by sampling method [62], low parasitism rates could be attributable to abiotic factors, such as climate. There is evidence that shows that aridity could be a driving force in reducing louse parasitism rates among birds inhabiting arid environments [11,15]. The area of South Texas where sampling occurred is a semi-arid habitat that can be heavily influenced by long periods of drought [63]. At the time of our study, South Texas was at the end of a three-year drought event, where the average precipitation during 2010–2012 was approximately 48.0 cm per year. This average is considerably lower than the average precipitation per year of 64.3 cm recorded from South Texas over a 30-year period (1981–2010 via [64]). Interestingly, when individual ranches were compared (San Antonio Viejo, which is arid and inland versus El Sauz, which is humid and coastal), there was no statistically significant difference in louse parasitism rates, which were 10.4% and 16.1%, respectively. It is therefore possible that during the time of this study El Sauz was not significantly more humid than San Antonio Viejo, possibly due to the drought (and despite different 30-year rainfall averages; see Methods). Notably, the intensity (or abundance) of lice on each host was higher on birds sampled from El Sauz than they were at San Antonio Viejo. For example, rather than finding one to three lice per host from San Antonio Viejo birds, birds from El Sauz (coastal climate, higher average rainfall) were generally parasitized by five or more lice per host individual, and in some instances as many as 30 lice on a single host. Therefore, it is a possibility that humidity does not have a substantial influence on louse prevalence and instead has more of an effect on intensity of individual host parasitism. A similar phenomenon has been described in previous studies in which host species at humid locations exhibit higher intensity of louse parasitism than arid locations [15,65].

With respect to phylogeny, within the amblyceran tree we recovered seven genetically distinct lineages, one of which we were able to morphologically identify (Figure 3). We discuss relationships among chewing louse lineages based primarily on the mitochondrial COI data set. We note for the most part that the nuclear data added additional support to the phylogeny. The only discrepancy was that the nuclear EF-1 α gene did not support a monophyletic Ischnocera. This has been documented in the literature several times (e.g., [41,45,56,66]), and is generally believed to be the result of poor taxonomic sampling. However, the fact that the slower evolving 18S did support a monophyletic Ischnocera implies that faster evolving EF-1 α may not be the best nuclear marker to use to reconstruct relationships at higher taxonomic levels [66].

Within the amblyceran tree (Figure 2), *Menacanthus* lice that parasitized Green Jay (*Cyanocorax yncas*) and Hooded Oriole (*Icterus cucullatus*) were almost identical genetically (average of 0.2% uncorrected *p*-distance) to the GenBank *Menacanthus* sp. from Bright-rumped Attila (*Attila spadiceus*); these likely represent the same species. It is unclear as to how these three behaviorally disparate taxa share this louse taxon; one possibility is transmission to wintering Hooded Orioles from tropical resident Bright-rumped Attila's, with subsequent transmission to Green Jays (which are sedentary) from breeding Hooded Orioles in Texas. Regardless, this *Menacanthus* subclade genetically differed on average 17.0% from the *Menacanthus* sp. parasitizing the Brown-crested Flycatcher (*Myiarchus*)

tyrannulus). Based on the amount of genetic divergence and a novel host association (Table 2), this *Myiarchus* flycatcher louse is likely a new species. A louse specimen we collected from a Northern Cardinal (Cardinalis cardinalis) was genetically similar to the louse parasitizing a Spangle-cheeked Tanager (Tangara dowii; 1.4% uncorrected p-distance). This could reflect the louse parasitizing the cardinal as being the same species (Menachanthus *eurysternus*) and it may have a large geographic range from South Texas to Costa Rica (and possibly much larger than that). The lice from Northern Parula (Setophaga americana) and Horned Lark (Eremophila alpestris) are on average 11.2% divergent from Menacanthus eurysternus (and 6% divergent from each other) and both represent novel host associations. The *Menacanthus eurysternus* complex is highly contentious; some believe it is widely distributed and parasitizes numerous host families and species [2,38], while others believe the complex contains subspecies or several similar species [67]. As such, its taxonomic status is uncertain [67]. Additionally, although an average of 11.2% genetic divergence is high, it may fall within the range of differentiation for a widespread species [9]. Thus, it is unclear if the Menacanthus lice parasitizing Setophaga americana and Eremophla alpetris represent a unique genetic lineage (even though both are new host associations) or normal diversity within a widespread louse lineage.

The genus *Myrsidea* is the most speciose genus in Phthiraptera, with 350 described species and new species frequently being described (e.g., [68–75]). In our study, there are six distinct genetic lineages within *Myrsidea* (Figure 2). High divergence was recorded in the louse collected from the Green Jay (*Cyanocorax yncas*) compared to its closest genetic relative (19.3%), a louse which parasitizes the Brown Jay (*Psilorhinus morio*) from Mexico. Notably, the Brown Jay has a similar distribution and belongs to the same family (Corvidae) as the Green Jay. Although we were unable to amplify any nuclear genes for the *Myrsidea* louse parasitizing the Green Jay, its COI sequence is genetically distinct and given this is a new host association (Table 2), this louse is likely new to science. A similarly high average divergence (18.7%) was recorded from the lice parasitizing the Tree Swallow (*Tachycineta bicolor*) as compared to the louse lineage parasitizing Carolina Wren (*Thryothorus ludovicianus*), Red-winged Blackbird (*Agelaius phoeniceus*), and Lark Sparrow (*Chondestes grammacus*). The *Myrsidea* found parasitizing these host taxa represent novel host associations, excluding the Carolina Wren (Table 2), and potentially new species.

Chewing lice found parasitizing the Louisiana waterthrush (Parkesia motacilla) have been previously recorded (identified as Myrsidea paleno [21]), but our study may be the first to obtain genetic data from this louse species (we did not have morphological material to definitively confirm this). There is a potential distinct Myrsidea lineage parasitizing Northern Cardinals (Cardinalis cardinalis) and the Black-throated Green Warbler (Setophaga virens), which represent two previously unrecorded host associations and were 12.6% divergent from other Myrsidea in this subclade. The lice collected from Northern Mockingbirds (Mimus polyglottos) are, on average, 14.4% genetically different from the closest GenBank sequence (Myrsidea nesomimi) collected from the Galapagos Mockingbird (Mimus parvulus). Although this genetic divergence is not quite as high as our 15% limit described above, it is still substantial and may reflect a new species especially when considering that little is known about Northern mockingbirds being parasitized by Myrsidea lice (Table 2). Notably, a previous phylogenetic study concluded that the Galapagos Mockingbird's closest relative is a member of the North American mockingbird genus, Mimus [76]. This analysis then led to the merging of the endemic Galapagos Mockingbird genus *Nesomimus* into *Mimus*. It is conceivable that when North American mockingbird species colonized the Galapagos Islands, they brought their parasites along with them, which subsequently diverged as their hosts diverged. This could potentially explain why the Myrsidea lice collected from Northern Mockingbirds in this study formed a sister clade to the lice from the Galapagos Mockingbird. The genus *Mimus* and the *Myrsidea* lice parasitizing them could therefore be a good study system for future studies of host-parasite coevolution.

Moreover, within *Myrsidea*, a Painted Bunting (*Passerina ciris*) was host to a distinct chewing louse lineage roughly 15.4% divergent from the sister clade of mockingbirds.

Given that this association is new to science (Table 2), it is likely that this Myrsidea lineage represents a new species. Additionally, the Myrsidea sp. parasitizing the Summer Tanager (Piranga rubra) is a novel host association (Table 2) and is 14.5% genetically different than its sister taxa (Myrsidea cruickshanki) that was found parasitizing Spangle-cheeked Tanager (Tangara dowii) and possibly represents a louse species new to science. Interestingly, the genus Piranga belonged to the same family as Tangara, Thraupidae, until 2003 when it was reclassified based on data showing that Piranga is genetically part of Cardinalidae [77]. A driving force for this split may be associated with geographic structuring as true thraupines includes species with a Neotropical distribution [77,78]. Perhaps the geographic structuring we see within the distantly related host species is being mirrored by the lice that parasitize them. Finally, within Myrsidea there is an interesting relationship with the louse Myrsidea textoris. The louse specimen of this species from GenBank was collected in South Africa from a Southern Masked Weaver (Ploceus velatus) and is known from several other South African host species including Lesser Masked Weaver (Ploceus intermedius) and Spectacled Weaver (Ploceus ocularis) [58]. In our study two separate hosts, Red-winged Blackbird and Brown-headed Cowbird (Molothrus ater), were likely parasitized by this louse species based on an average 4.9% genetic divergence (uncorrected *p*-distance). Another *Ploceus* host species, African Village Weaver (Ploceus cucullatus) was introduced to Hispanolia [79] and is a known host of Myrsidea textoris. These established Hispanolia populations of Village Weavers have been recorded flocking with Shiny Cowbirds (Molothrus bonariensis) in the Dominican Republic [80]. Due to proximity, transmission from Village Weavers to Shiny Cowbirds could have occurred. The Shiny Cowbird has distributional overlap with both the Red-winged Blackbird and Brown-headed Cowbird in southern Florida and parts of Central America. These past introductions and distribution overlaps of host species could possibly explain the intercontinental distribution for Myrsidea textoris.

Ricinus lice formed several sub-clades (Figure 2). One of these was Ricinus parasitizing the Golden-winged Warbler (Vermivora chrysoptera), Tennessee Warbler (Oreothlypis peregrina), Northern Waterthrush (Parkesia noveboracensis), and Bay-breasted Warbler (Setophaga castanea) which formed one sub-clade, are on average 12.2% genetically different from each other, and this sub-clade is sister to Ricinus mugimaki parasitizing a host from South Africa (Chorister Robin-chat, Cossypha dichroa). When looking at known louse-host associations, different species of *Ricinus* are known to parasitize each of the host species within this clade. There are known host associations of *Ricinus picturatus* parasitizing members from the genus Vermivora [81] such as the Golden-winged Warbler and the former Vermivora member (now Leiothlypis), Tennessee Warbler. There are records of Ricinus seiuri parasitizing the Northern Waterthrush and the Bay-breasted Warbler has a known association with Ricinus dendroicae. Unfortunately, there are no molecular data available for these particular Ricinus species. Given this lack of genetic information coupled with the inability to morphological identify these lice to species, we can only discern that there are potentially three louse species in this clade based on genetic divergence and host associations. A last subclade reflects Ricinus sp. parasitizing a Varied Bunting (Passerina versicolor) and a Vermilion Flycatcher (*Pyrocephalus rubinus*), which are genetically almost identical to a *Ricinus* sp. parasitizing (Cyanocompsa parellina).

Finally, we found in the amblyceran tree a single species of *Hohorstiella (passerinae)* parasitizing a Common Ground Dove (*Columbina passerinae*); this species is known to parasitize other dove species to include Inca Dove (*Columbina inca*; Figure 2).

Within the ischnoceran tree, we recovered 14 genetically distinct lineages, several of which we were able to morphologically identify (Figure 3). This is not particularly surprising as numerous ischnoceran species and host associations are being discovered and described globally (e.g., [58,82–88]). However, these new descriptions often come from areas that are traditionally understudied in terms of host and louse diversity. The fact that there are 14 distinct lineages from South Texas is somewhat surprising; nine of these come from one genus and four are known louse–host associations that lack genetic reference material.

All Brueelia in this study were genetically identical or nearly identical to sequences obtained from GenBank, with the exception of two specimens (Figure 3). The two exceptions were collected from a Verdin (Auriparus flaviceps), which has a known association with Brueelia audax and Bell's Vireo (Vireo bellii), and they are an average of 13.0% and 14.1% different from other Brueelia in their subclade, respectively. These are potential unique genetic lineages coupled with novel host associations (Table 2), revealing a strong possibility of new species within Brueelia. There have been numerous studies over the past few years that have focused efforts on collecting Brueelia sp. and using molecular analysis to resolve the Brueelia complex [89]. Recently Bush et al. [90] recognized this group as being paraphyletic and Gustafsson and Bush [91] proposed a revised classification of the complex based on morphological data and excluding host relationships. This genus is one of the largest groups within Ischnocera, originally containing over 260 described species of which roughly 90% are host specific [92] and further investigation increased this number to 426 species [91]. In fact, our phylogeny reflects specificity for some species to host family (Figure 3). For example, one Brueelia lineage was found to only parasitize species located in the bird family Mimidae (Figure 3). This lineage likely represents individuals of Bruellia brunneinucha based on minimal genetic divergence from the GenBank specimen and because this louse species targets other mimids as well, with described associations with Tropical Mockingbird (Minus gilvus), Blue-and-white Mockingbird (Melanotis hypoleucus), and Bahama Mockingbird (Mimus gundlachii [93]). Another example of host family specificity applies to Bruellia xanthocephali and Bruellia ornatissima, which collectively form a clade that appears to only parasitize members of family Icteridae (Figure 3). However, there is also a group of hosts that appears to be parasitized by a non-host specific louse, Bruellia dorsale; this louse is recovered as parasitizing four different host species from three different families (Figure 3). Interestingly, the host species that form that clade frequently inhabit thorn-scrub and dense vegetation. This suggests that Bruellia dorsale may represent an opportunistic parasite within that particular habitat. Our results for Brueelia support the conclusions of Johnson et al. [92], who regarded Brueelia as having members that were both host specific and host generalists.

The *Penenirmus* collected from the Golden-fronted Woodpecker (*Melanerpes aurifrons*) was highly supported as sister to a louse collected from another Picidae species, the African Gray Woodpecker (*Chloropicus goertae*) although these two louse species were highly divergent (21.3%). Sister to the clade of *Penenirmus* woodpecker lice are two other louse genera, *Lunaceps* and *Cummingsiella*, to the exclusion of the *Penenirmus* parasitizing the Bushtit (*Psaltriparus minimus*). This renders the genus *Penenirmus* polyphyletic, where different lineages parasitize Piciformes and Passeriformes hosts. These results, although representing limited sampling, support previous literature finding a lack of monophyly for *Penenirmus* [45,58,94] and the necessity for taxonomic revision of this louse genus.

While the genus *Philopterus* is well supported, it is effectively a large polytomy with high amounts of divergence (average uncorrected *p*-distance 20.7%) among lineages (Figure 3). As such, it is difficult to determine relationships within this genus. This difficulty is exacerbated by the general lack of genetic references on GenBank until recently [84]. Unfortunately, many of the samples in this clade were nymphs, thus it was difficult for us to identify any of these samples to species. Of the nine recorded distinct lineages within this genus (22.3% average *p*-distance), seven are associated with novel host associations (Figure 3, Table 2). Most of the records detailing *Philopterus* and their host associations in North America are many decades old (e.g., [95,96]) and are based solely on morphology. In general, there is little recent information providing molecular data or new host associations about the genus apart from Price et al. [1], Mey [97], Bush et al. [11], and Najer et al. [84]. Clearly, additional molecular and morphological studies focused on *Philopterus* are needed.

A well supported clade that is referred to as the *Degeeriella* complex by Clay [98] was recovered within the ischnoceran phylogeny (Figure 3). This clade, which comprises several genera including *Degeeriella*, *Picicola*, *Trogoninirmus*, and *Cotingacola*, is often defined as being strongly supported based on morphology and molecular data, although

many genera within this group have been shown to be paraphyletic [99,100]. Specifically, the genus *Picicola* has been shown to be paraphyletic in many studies (e.g., [28,45,100]), including our own (Figure 3). Looking into the complex a bit further, we have a distinct lineage parasitizing the Scissor-tailed flycatcher (*Tyrannus forficatus*) that is an average of 22.4% divergent from the clade containing the *Picicola* lice parasitizing *Melanerpes* species. While *Picicola foedus* is known to parasitize the Scissor-tailed flycatcher, molecular data demonstrating this association are not available. Therefore, we provide new molecular information about this host association. If the louse from this study is found to be morphologically different from the louse identified in the original louse–host association, this could potentially represent a new species.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/ 10.3390/d13090430/s1. Figure S1: Bayesian phylogeny of nuclear gene (18S). Figure S2: Bayesian phylogeny of nuclear gene (EF-1 α). Figure S3: Bayesian phylogeny of combined nuclear genes. Figure S4: Bayesian phylogeny of combined mitochondrial (COI gene) and nuclear gene (18S). Figure S5: Bayesian phylogeny of combined mitochondrial (COI gene) and nuclear gene (EF-1 α). Table S1: Louse sequences obtained from GenBank. Table S2: Louse specimens used in analysis from South Texas. Table S3: Avian species sampled for lice in south Texas.

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